## Claims:

- An isolated nucleic acid molecule encoding a hyperimmune serum reactive antigen or a fragment thereof comprising a nucleic acid sequence which is selected from the group consisting of:
  - a) a nucleic acid molecule having at least 70% sequence identity to a nucleic acid molecule selected from Seq ID No 3-4, 16, 19-21, 28-29, 33-38, 41-42, 44, 48-52, 55, 57-58, 61, 63, 65, 67-68, 72, 74-75, 81, 84, 91, 94, 96-97, 101, 105-108, 112, 115-117, 119, 123-178,
  - b) a nucleic acid molecule which is complementary to the nucleic acid molecule of a),
  - c) a nucleic acid molecule comprising at least 15 sequential bases of the nucleic acid molecule of a) or b)
  - d) a nucleic acid molecule which anneals under stringent hybridisation conditions to the nucleic acid molecule of a), b), or c)
  - e) a nucleic acid molecule which, but for the degeneracy of the genetic code, would hybridise to the nucleic acid molecule defined in a), b), c) or d).
- 2. The isolated nucleic acid molecule according to claim 1, wherein the sequence identity is at least 80%, preferably at least 95%, especially 100%.
- 3. An isolated nucleic acid molecule encoding a hyperimmune serum reactive antigen or a fragment thereof comprising a nucleic acid sequence selected from the group consisting of
  - a) a nucleic acid molecule having at least 96% sequence identity to a nucleic acid molecule selected from Seq ID No 8-10, 13-15, 17-18, 24, 27, 32, 39-40, 45-47, 56, 59, 62, 69-70, 73, 77, 79, 82, 85-86, 88, 90, 103, 109-110, 114, 121.
  - b) a nucleic acid molecule which is complementary to the nucleic acid molecule of a),
  - c) a nucleic acid molecule comprising at least 15 sequential bases of the nucleic acid molecule of a) or b)
  - d) a nucleic acid molecule which anneals under stringent hybridisation conditions to the nucleic acid molecule of a), b) or c),
  - e) a nucleic acid molecule which, but for the degeneracy of the genetic code, would hybridise to the nucleic acid defined in a), b), c) or d).
- 4. An isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of
  - a) a nucleic acid molecule selected from Seq ID No 5, 7, 30-31, 53, 60, 66, 76, 83, 87, 92, 99, 120,
  - b) a nucleic acid molecule which is complementary to the nucleic acid of a),
  - c) a nucleic acid molecule which, but for the degeneracy of the genetic code, would hybridise to the nucleic acid defined in a), b), c) or d).
- 5. The nucleic acid molecule according to any one of the claims 1, 2, 3 or 4, wherein the nucleic acid is DNA.
- 6. The nucleic acid molecule according to any one of the claims 1,2, 3, 4, or 5 wherein the nucleic acid is RNA.
- 7. An isolated nucleic acid molecule according to any one of claims 1 to 5, wherein the nucleic acid molecule is isolated from a genomic DNA, especially from a *H. pylori* genomic DNA.
- 8. A vector comprising a nucleic acid molecule according to any one of claims 1 to 7.
- A vector according to claim 8, wherein the vector is adapted for recombinant expression of the hyperimmune serum reactive antigens or fragment thereof encoded by the nucleic acid molecule according to any one of claims 1 to 7.

- A host cell comprising the vector according to claim 8 or 9.
- 11. A hyperimmune serum-reactive antigen comprising an amino acid sequence being encoded by a nucleic acid molecule according to any one of the claims 1, 2, 5, 6 or 7 and fragments thereof, wherein the amino acid sequence is selected from the group consisting of Seq ID No 181-182, 194, 197-199, 206-207, 211-216, 219-220, 222, 226-230, 233, 235-236, 239, 241, 243, 245-246, 250, 252-253, 259, 262, 269, 272, 274-275, 279, 283-286, 290, 293-295, 297, 301-356.
- 12. A hyperimmune serum-reactive antigen comprising an amino acid sequence being encoded by a nucleic acid molecule according to any one of the claims 3, 5, 6, or 7 and fragments thereof, wherein the amino acid suequece is selected from the group consisting of Seq ID No 186-188, 191-193, 195-196, 202, 205, 210, 217-218, 223-225, 234, 237, 240, 247-248, 251, 255, 257, 260, 263-264, 266, 268, 281, 287-288, 292, 299.
- 13. A hyperimmune serum-reactive antigen comprising an amino acid sequence being encoded by a nucleic acid molecule according to any one of the claims 4, 5, 6, or 7 and fragments thereof, wherein the amino acid sequence is selected from the group consisting of **Seq ID No** 183, 185, 208-209, 231, 238, 244, 254, 261, 265, 270, 277, 298.
- Fragments of hyperimmune serum-reactive antigens selected from the group consisting of peptides 14. comprising amino acid sequences of column "predicted immunogenic aa" and "location of identified immunogenic region" of Table 1, the serum reactive epitope of Table 3 especially peptides comprising amino acid 63-91, 95-101, 110-116, 134-148, 150-156, 158-164, 188-193, 197-209, 226-241, 247-254, 291-297, 312-319, 338-346, 351-358, 366-378, 404-410, 420-438, 448-454, 465-473, 482-488, 490-498, 503-510, 512-519, 531-543, 547-554, 568-575, 589-604, 610-631 and 239-308 of Seq ID No 179; 16-29, 35-47, 50-68, 70-79, 91-101, 143-149, 158-163, 185-191, 196-206, 215-224, 230-237, 244-251, 258-278, 290-311, 319-325, 338-351, 365-385, 396-429, 445-454, 458-466, 491-499, 501-521, 17-79 and 218-233 of Seq ID No 180; 4-10, 16-41, 46-66, 77-84, 91-97, 102-118, 125-144, 187-200, 202-214, 245-253, 255-261, 286-295, 300-330, 335-342, 350-361, 363-381, 385-392, 396-416, 435-450 and 460-470 of Seq ID No 181; 11-19, 27-48, 52-59, 77-82, 84-107, 118-125, 127-154, 178-183, 192-209, 215-221, 286-295, 302-313, 350-357, 402-415, 417-431, 453-463, 465-493 and 313-331 of Seq ID No 182; 19-26, 30-43, 47-55, 63-68, 72-80, 97-104, 107-119, 129-146, 160-175, 194-216, 231-251, 254-260 and 26-43 of Seq ID No 183; 7-13, 29-37, 65-81, 110-120, 123-131, 135-152, 230-249, 254-260, 284-290, 292-299, 317-326, 329-336, 403-444, 452-458, 466-477, 490-498, 510-519, 541-550, 557-566 and 533-567 of Seq ID No 184; 5-47, 71-77, 79-86, 89-95, 120-126, 137-144, 176-181, 184-196, 202-208, 211-232, 236-282, 301-313, 317-325, 341-347, 353-384, 394-400, 412-433, 436-443 and 59-75 of Seq ID No 185; 4-18, 22-38, 59-69, 106-112, 116-130, 138-149, 156-170, 175-197, 200-214, 216-223, 233-244, 255-261, 266-276, 279-286, 325-333, 342-348, 366-399, 402-420, 429-441, 1-104 and 130-147 of Seq ID No 186; 50-58, 69-95, 97-113, 131-136, 157-163, 170-175, 188-212, 220-226, 254-259, 265-277, 283-289, 297-308, 311-318, 347-358, 360-369, 378-401, 416-421, 440-450, 454-462, 470-476, 493-502, 506-514, 536-567, 585-590, 598-607, 613-618, 653-659 and 35-46 of Seq ID No 187; 16-29, 32-60, 65-87, 89-123, 128-134, 137-158, 162-173, 178-196, 210-216, 218-228 and 206-225 of Seq ID No 188; 10-20, 26-35, 51-64, 86-91, 94-100, 113-122, 154-160, 185-191, 193-201, 211-217, 225-230, 237-246, 251-257, 298-304, 306-312, 316-328, 340-348, 357-389, 391-397, 415-421, 449-456, 458-471, 488-495, 502-511, 24-55 and 236-341 of Seq ID No 189; 5-22, 41-51, 87-93, 114-122, 127-136, 150-156, 158-166, 223-233, 245-263, 291-296, 9-126 and 127-285 of Seq ID No 190; 30-43, 46-56, 61-70, 72-83, 85-93, 103-113, 119-125, 151-166, 179-191, 212-218, 225-231, 236-243, 262-267, 291-307, 331-344, 349-355, 366-372, 380-386, 414-422, 428-447, 459-464, 469-478, 507-519, 525-544, 563-569, 576-590, 620-626, 633-643, 654-659, 665-671, 684-707, 717-723, 725-733, 747-779, 782-801 and 347-361 of Seq ID No 191; 4-12, 14-26, 37-80, 107-115, 133-139, 144-150, 154-165, 173-180, 191-199, 205-211, 221-231, 237-244, 254-284, 307-340, 342-353, 360-368, 370-380, 479-493, 495-503,

- 84 -

509-522, 525-536, 539-547, 554-560, 565-573, 578-583, 7-23 and 465-479 of Seq ID No 192; 4-17, 47-55, 76-83, 85-100, 104-112, 117-123, 126-135, 142-148, 156-167, 174-182, 267-273 and 258-283 of Seq ID No 193; 8-32, 36-42, 65-88, 102-108, 112-140, 147-163, 170-179, 183-193 and 117-124 of Seq ID No 194; 12-18, 45-50, 62-77, 82-95, 99-113, 115-123, 125-147, 155-177, 187-209, 211-223, 244-253, 259-270, 278-297, 302-307, 311-318, 329-334, 350-356, 359-365, 390-400, 402-413 and 333-350 of Seq ID No 195; 4-13, 15-27, 30-46, 53-58, 68-74, 82-95, 115-126, 134-139, 148-153, 159-176, 182-199, 201-217, 220-225, 227-235, 237-248, 253-266, 300-315, 322-336, 390-396, 412-426, 438-445, 448-459, 477-484, 502-508, 515-527, 529-537, 553-568, 643-651, 658-667, 690-703 and 376-400 of Seq ID No 196; 4-10, 24-32, 38-55, 59-67, 70-77, 80-87, 89-97, 123-129, 134-151, 166-172, 178-189, 191-216, 218-235, 245-259, 271-315, 326-339, 341-360 and 73-94 of Seq ID No 197; 13-25, 31-38, 43-57, 79-85, 92-99, 106-112, 117-128, 130-139, 146-158, 160-175, 194-204, 211-222, 225-232, 234-242, 263-270, 278-292, 299-320, 322-333 and 240-256 of Seq ID No 198; 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4-16, 37-59, 64-70, 79-87, 93-102, 107-127, 143-165, 172-188, 197-204, 207-218, 221-227, 242-248, 258-277, 289-296, 298-316, 332-338, 344-365, 367-373, 375-382, 400-408, 415-425, 438-446 and 235-250 of Seq ID No 205; 4-37, 39-66, 84-98, 101-127, 140-149, 157-163, 166-172, 175-182, 184-193, 203-208, 215-232, 234-247, 250-299, 303-345 and 183-204 of Seq ID No 206; 10-20, 41-61, 73-87, 112-141, 176-192, 194-201, 205-222, 230-237, 257-264, 276-282, 284-310, 312-318, 330-337, 349-357 and 304-328 of Seq ID No 207; 4-31, 42-103, 105-113, 121-153, 160-181, 188-196, 210-226, 231-264, 272-287, 297-304, 328-336 and 304-318 of Seq ID No 208; 21-43, 46-52, 54-70, 72-79, 94-107, 133-141, 160-166, 217-253, 311-317, 359-365, 374-381, 390-395, 434-440, 488-494, 497-502, 511-522, 554-563, 565-574, 577-585, 591-598, 601-606, 617-625, 633-643, 658-664, 676-682, 694-702, 710-719, 754-760, 782-788, 802-808, 916-921, 942-948, 955-964, 973-979, 992-998, 1006-1011, 1016-1023, 1030-1038, 1046-1053, 1059-1066, 1088-1098, 1119-1126, 1129-1135, 1156-1171, 1173-1181, 1202-1210, 1255-1261, 1268-1280, 1295-1310, 1312-1320, 1375-1381, 1406-1417, 1450-1471, 1478-1492, 1498-1506, 1569-1578, 1603-1608, 1611-1624, 1648-1655, 1663-1670, 1680-1698, 1702-1707, 1713-1719, 1737-1742, 1747-1753, 1762-1769, 1771-1785, 1790-1804, 1811-1818, 1830-1836, 1838-1852, 1874-1886, 1893-1899, 1902-1909, 1942-1948, 1952-1962, 1980-1986, 2001-2017, 2020-2028, 2042-2050, 2052-2068, 2074-2079, 2083-2095, 2107-2113, 2147-2155, 2177-2194, 2203-2211, 2236-2241, 2251-2258, 2267-2274, 2285-2292, 2314-2328, 2330-2340, 2358-2365, 2390-2401, 2408-2418, 2432-2453, 2463-2476, 2486-2507, 2528-2537, 2540-2548, 2552-2558, 2568-2576, 2596-2601, 2610-2622, 2629-2638, 2653-2669, 2718-2727, 2749-2767, 2777-2784, 2789-2795, 2806-2815, 2817-2824, 2835-2843, 2847-2854, 2860-2881, 511-523, 612-630 and 1790-1803 of Seq ID No 209; 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64, 86-94, 97-104, 121-129, 257-272, 279-286, 288-294, 307-327, 334-340, 369-375, 377-386, 406-412. 418-423, 430-438, 441-447, 459-465, 469-476, 482-488, 510-546, 550-580, 584-622, 638-645, 653-659, 675-683, 692-705, 723-731, 752-761, 788-795 and 54-72 of Seq ID No 215; 11-33, 36-46, 88-104, 116-126, 134-170, 189-195, 199-217, 225-250, 255-261, 266-273, 280-291, 296-313, 334-341, 343-349, 354-360, 362-369, 373-380, 387-401, 406-420 and 259-273 of Seq ID No 216; 9-14, 28-44, 57-64, 72-79, 86-93, 104-111, 116-126, 142-150, 159-164 and 61-86 of Seg ID No 217; 10-17, 26-33, 43-61, 69-95, 101-107, 109-125, 129-135, 137-144, 147-153, 158-169, 177-187, 209-219, 221-232, 235-247, 261-268, 271-282, 296-302, 306-347, 355-362, 364-379, 386-399, 409-418, 424-442, 451-460, 467-479, 490-498 and 60-74 of Seq ID No 218; 8-14, 20-31, 65-84, 94-99, 154-179, 193-207, 238-253 and 96-118 of Seq ID No 219; 4-24, 30-44, 47-62, 84-93, 108-116, 124-133, 136-141, 201-209, 217-223, 228-235, 238-245, 247-270, 275-285, 290-314, 328-338, 342-349, 353-365, 375-383, 386-392, 394-402, 417-427, 443-459, 465-481, 492-514, 516-524, 550-566, 602-617, 630-639, 666-676, 687-693, 719-730, 747-753, 783-790, 799-816, 824-831, 837-842 and 167-189 of Seq ID No 220; 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4-21, 64-71, 73-84, 128-138, 144-162, 203-217, 240-263, 288-298, 300-308, 310-317, 325-351, 369-380, 391-411 and 330-345 of Seq ID No 233; 5-11, 25-31, 39-48, 51-79, 89-98, 100-122, 135-148, 166-201, 203-227, 230-250, 254-260, 266-272, 274-282, 299-305, 328-337 and 31-45 of Seq ID No 234; 12-23, 29-48, 51-60, 66-72, 75-81, 83-93, 103-115, 133-148, 168-174, 195-204, 222-229, 231-240, 242-251, 270-280, 286-305, 322-344, 349-360, 364-370, 378-400, 421-441, 448-484, 486-493, 495-501, 504-534, 547-561, 567-590, 597-607, 621-635, 643-649, 658-685, 688-694, 702-711, 717-731, 737-742, 759-765, 767-772, 776-786, 803-809, 815-825, 854-908, 910-919, 923-930, 942-948, 961-975, 994-1014 and 915-940 of Seq ID No 235; 4-9, 32-47, 51-61, 75-96, 139-191 and 1-124 of Seq ID No 236; 4-13, 17-38, 43-49, 55-76, 88-95, 110-121, 128-146, 151-157, 162-214, 222-240, 243-249, 251-273, 275-281, 292-298, 300-309, 312-320, 322-331, 355-369, 376-408, 446-460, 471-482, 485-509

and 191-203 of Seq ID No 237; 4-21, 72-82, 89-103, 106-115, 118-124, 140-146, 174-184, 191-200, 204-213, 218-224, 261-266, 282-293, 299-309, 311-340, 342-358, 362-372, 381-389, 391-402, 413-421, 438-447, 457-464, 470-478, 501-507, 545-560, 578-624, 631-641, 658-670, 680-689, 717-738, 753-759, 795-805. 816-822, 830-838, 842-848, 869-881, 892-898, 33-51 and 818-835 of Seq ID No 238; 4-21, 79-85, 156-177, 183-188, 206-214, 243-249, 261-269, 287-292, 315-322, 334-345, 360-366, 374-390, 402-411, 37-97 and 260-399 of Seg ID No 239; 4-9, 19-54, 58-78, 97-104, 111-120, 126-134, 137-145, 163-173, 178-188, 193-203, 211-224, 246-286, 288-324, 337-346, 355-362, 374-390, 392-398, 409-417 and 240-249 of Seq ID No 240; 5-12, 14-31, 35-41, 43-61, 82-92, 97-105, 134-145, 155-166, 184-203, 215-223, 225-251, 272-279, 281-306, 310-345, 358-418, 435-473, 482-490, 525-532, 538-547, 549-563, 578-604, 613-639 and 144-154 of Seq ID No 241; 53-59, 64-72, 74-100, 133-152, 154-172, 176-181, 207-214, 225-238, 275-297, 304-310, 331-340, 362-367, 384-395, 403-410, 437-443, 448-456, 482-490, 579-597, 602-610, 625-630, 633-651, 699-707, 709-715, 734-743, 750-762 and 544-685 of Seq ID No 242; 12-18, 22-40, 45-83, 89-97, 103-109, 147-153, 159-173, 195-204, 210-219, 243-253, 259-265, 273-282, 303-309, 315-325, 332-340, 346-358, 362-367, 377-390, 393-402, 418-426, 447-455, 467-480, 505-512, 514-525, 548-561, 566-576, 584-596, 619-626, 638-645, 649-659, 661-680, 699-708, 714-720, 753-759, 766-772, 775-781, 801-808, 202-218, 282-299, 339-350 and 617-628 of Seq ID No 243; 5-33, 52-62, 87-101, 111-135, 137-143, 145-152, 190-202, 209-221, 233-245, 253-270 and 151-215 of Seq ID No 244; 19-29, 32-39, 42-48, 75-94, 124-135, 137-145, 152-160, 176-182, 193-203, 215-236, 266-273, 275-291, 297-306, 311-319, 322-342, 348-360, 369-378, 394-401 and 48-64 of Seq ID No 245; 4-11, 13-33, 36-43, 53-63, 65-80; 112-129, 134-141, 143-155, 157-168, 178-188, 191-199, 201-207, 215-229, 242-255, 263-270, 283-315, 320-329, 333-338, 340-349, 412-426, 465-478, 485-490, 498-512, 540-554 and 390-516 of Seq ID No 246; 4-18, 23-32, 41-47, 54-70, 88-99, 104-111, 118-138, 143-148, 150-162, 168-175, 181-188, 203-211, 214-220, 227-245, 251-268, 275-281, 287-296, 323-333 and 1-90 of Seq ID No 247; 8-34, 38-49, 72-83, 85-91, 94-104, 112-125, 134-142, 148-168, 181-189, 191-198, 202-214, 222-233, 242-254, 256-262, 273-278, 287-294, 314-325 and 141-159 of Seq ID No 248; 4-24, 30-36, 47-75, 82-105, 124-134, 151-157, 192-202, 208-214, 219-226, 234-247, 285-290, 318-324, 332-340, 343-349, 380-386, 453-462, 472-478, 484-501, 531-540, 550-557, 604-612, 620-625, 642-648, 652-671, 64-84, 93-180 and 181-446 of Seq ID No 249; 12-18, 24-32, 68-75, 77-83, 96-101, 109-116, 129-136, 152-164, 175-184, 190-199, 206-215, 224-233, 241-250, 258-264, 273-292, 302-312, 319-331, 334-346, 348-368, 387-395, 408-416, 420-429, 437-452 and 364-374 of Seq ID No 250; 11-28, 36-52, 60-67, 74-79, 108-116 and 61-76 of Seq ID No 251; 20-27, 38-49, 69-74, 84-107, 138-145, 161-168, 179-195, 210-226, 228-252, 267-281, 283-296, 305-311, 333-340, 342-356, 361-372, 380-399, 401-414, 458-466, 475-481, 492-507, 515-520 and 146-160 of Seq ID No 252; 43-61, 68-74, 76-90, 120-128, 130-149, 156-161, 164-182, 206-234, 242-252, 269-274, 291-304, 332-345, 349-355, 360-371, 374-388, 434-440, 447-453, 459-465, 469-496, 504-522 and 261-285 of Seq ID No 253; 4-17, 24-30, 37-49, 87-98, 118-124, 126-136, 144-171, 176-188, 206-214, 216-228, 233-240, 246-252, 262-271, 277-297, 307-330, 333-342, 346-352, 355-361, 368-386, 391-400, 413-420, 474-480 and 401-427 of Seq ID No 254; 15-26, 31-46, 51-72, 80-93, 96-109, 131-137, 150-158, 179-185, 189-209, 211-219, 221-234, 241-247, 255-262, 265-271, 283-288 and 173-190 of Seq ID No 255; 28-37, 39-45, 51-58, 77-84, 89-97, 132-148, 171-180, 199-205, 212-218, 220-226, 257-265, 273-300, 307-327, 334-340, 344-365, 385-390, 402-408, 426-436, 450-468, 476-485 and 425-497 of Seq ID No 256; 4-25, 70-76, 80-88, 90-100, 120-128, 162-169, 183-203, 261-277, 279-289, 291-297, 302-308, 321-327, 339-353, 358-377, 392-401, 404-410, 414-422, 443-450, 456-461, 470-488, 490-497, 510-535, 570-611, 618-630, 639-647, 649-660, 668-690, 702-716, 718-724, 737-747, 750-764 and 497-509 of Seq ID No 257; 12-48, 50-64, 99-108, 216-223, 235-241, 244-254, 262-274, 287-293, 310-316, 320-326, 361-366, 377-383, 390-395, 408-414, 418-425, 438-444, 462-469, 494-505, 524-530, 536-547, 551-566, 592-598, 601-613, 678-685, 687-695, 709-717, 727-737, 751-757, 760-765, 772-778, 782-788, 801-807, 822-830, 859-868, 870-878, 884-890, 898-903, 909-919, 953-969, 973-980, 990-1000, 1002-1019, 1041-1047, 1059-1065, 1090-1095, 1116-1127, 1130-1139, 1143-1149, 1151-1168, 1178-1183, 1188-1195, 1197-1209, 1213-1220, 1226-1234, 1236-1247, 1255-1274, 1276-1282, 76-100, 270-284, 309-438, 493-505, 786-942 and 947-967 of Seq ID No 258; 4-9, 24-34, 46-95, 97-109, 119-130 and 138-156 of Seg ID No 259; 9-26, 28-35, 43-53, 55-68, 83-92, 99-105, 110-135, 139-149, 157-162, 164-170, 173-183, 193-208, 210-230, 239-245, 253-259, 263-271, 293-305, 310-320, 322-331, 336-343, 351-364, 367-376, 92-107 and 154-173 of Seq ID No 260; 19-39, 52-62, 108-117, 145-152, 160-168, 194-203, 229-240, 252268, 280-287, 308-316, 333-339, 383-390, 403-412, 414-424, 438-445, 464-472, 479-484, 489-505, 510-526 and 247-260 of Seq ID No 261; 5-17, 25-52, 60-77, 105-113, 118-125, 162-167, 228-234, 272-279, 328-334, 341-357, 381-395, 400-406, 512-518, 557-569, 586-592, 645-651, 690-695, 701-709, 720-726, 733-743, 751-758, 781-786, 879-886, 929-934, 939-944, 952-960, 965-975, 994-1001, 1039-1045, 1102-1109, 1164-1181, 1198-1206, 1223-1229, 1253-1259, 1283-1292, 1312-1317, 1339-1349, 1360-1370, 1389-1398, 1400-1412, 1452-1465, 1470-1484, 1490-1497, 1519-1525, 1554-1564, 1578-1591, 1623-1636, 1638-1646, 1669-1679, 1685-1697, 1704-1711, 1713-1720, 1730-1736, 1738-1749, 1756-1764, 1778-1786, 1796-1803, 1817-1826, 1849-1866, 1975-1993, 2017-2032, 2044-2053, 2070-2086, 2091-2109, 2116-2127, 2156-2167, 2182-2188, 2197-2202, 2244-2252, 2281-2287, 2290-2307, 2350-2361, 2383-2404, 2425-2433, 2445-2455, 2495-2505 and 394-549 of Seq ID No 262; 9-24, 31-53, 57-67, 69-79, 84-114, 133-141, 144-172, 178-186 and 13-46 of Seq ID No 263; 4-25, 27-35, 43-52, 59-70, 79-91, 115-130, 136-152, 154-163, 170-179 and 1-58 of Seq ID No 264; 4-30, 49-55, 71-80, 96-105, 111-126, 139-146, 149-162, 239-245, 279-285, 290-296, 300-307, 331-337, 343-350 and 250-351 of Seq ID No 265; 9-27, 34-41, 43-51, 92-111, 114-120, 123-131, 139-150, 156-171, 176-186, 188-204, 229-241, 252-258, 266-279, 288-297, 319-334, 338-348, 373-379, 389-398, 431-439, 479-484 and 214-398 of Seq ID No 266; 4-15, 18-27, 47-52, 68-83, 91-97, 104-110, 115-121, 139-147, 157-164, 198-206, 227-236, 241-254, 264-273, 278-289, 311-320, 353-361, 372-383, 405-420, 426-434 and 232-386 of Seq ID No 267; 4-10, 24-34, 91-97, 129-141, 156-163, 184-190, 205-219, 229-235, 256-273, 278-285 and 93-116 of Seq ID No 268; 7-29, 35-54, 71-83, 85-91, 104-111, 122-134, 138-144, 146-154, 158-174, 177-183, 186-201, 207-215, 223-235, 240-247, 262-273, 275-283, 287-292 and 48-66 of Seq ID No 269; 7-27, 31-47, 49-70, 75-102, 110-149, 157-171, 217-223, 235-251, 294-302, 358-364, 367-375, 387-393, 395-412, 423-430, 441-451, 456-470, 472-486, 488-495, 499-509, 515-529, 536-549, 556-570, 574-603, 607-615, 625-633, 642-658, 670-676, 683-702, 708-716, 720-726, 747-756, 763-784, 803-812, 815-826 and 475-490 of Seq ID No 270; 7-22, 30-38, 53-59, 64-75, 83-95, 97-112, 120-131, 133-142, 145-151, 154-166, 172-180, 189-203, 227-238, 277-287, 9-156 and 174-287 of Seq ID No 271; 13-23, 25-32, 111-117, 150-164, 185-193, 207-212, 216-224, 230-236, 263-272, 304-311, 342-348, 374-385, 391-407, 444-458, 480-487, 489-499, 523-542, 544-558, 572-579, 620-640, 686-696, 703-710, 742-755, 765-772, 817-822, 830-837, 865-872, 931-937 and 66-86 of Seq ID No 272; 4-27, 49-56, 62-70, 86-92, 121-127, 151-163, 170-182, 195-202, 212-226, 237-243 and 234-254 of Seq ID No 273; 4-10, 13-24, 39-51, 62-78, 92-104, 107-117, 134-141, 156-161, 166-181, 210-216, 222-229, 256-266, 273-280, 297-304, 313-330, 336-349, 371-376, 433-439, 443-448, 488-493, 506-515, 527-534, 560-572, 575-583, 587-593 and 252-483 of Seq ID No 274; 4-15, 21-38, 45-56, 81-95, 102-108, 118-130, 133-147, 152-162, 166-171, 199-204, 211-218, 230-240, 253-261, 274-283, 288-294, 312-317, 325-336, 344-357, 391-414 and 24-146 of Seq ID No 275; 26-31, 38-56, 65-82, 90-101, 112-119, 123-153, 175-188, 197-216, 234-242, 249-265, 273-286, 290-305, 327-335, 338-346, 361-372, 394-404 and 290-306 of Seq ID No 276; 17-26, 43-48, 50-73, 81-93, 95-107, 139-146, 158-168, 171-176, 190-196, 202-212, 216-223, 243-266, 274-282, 308-313, 324-330, 344-378, 380-387, 403-422, 427-443, 448-455, 457-465, 491-515, 517-528, 553-567, 589-599, 610-617, 642-648, 670-697, 709-717, 726-743, 745-759, 769-803, 807-823, 840-849 and 820-851 of Seq ID No 277; 4-18, 39-48, 53-63, 66-90, 102-117, 125-134, 137-145, 156-162, 169-197, 26-40 and 56-80 of Seq ID No 278; 21-33, 36-42, 49-60, 68-76, 91-105, 123-130, 141-161, 169-178, 185-190, 192-199, 205-214, 223-233, 239-247, 260-269, 284-293, 300-314, 324-352, 357-364, 373-382, 389-403, 420-432, 438-446, 466-471, 477-484, 503-509, 549-556, 558-576, 600-623, 625-635, 654-661, 663-669, 671-687, 702-716, 735-741, 744-750, 757-766, 776-786, 807-815, 824-832, 854-860, 863-897, 909-915, 920-946, 952-959, 982-997, 1024-1038, 1049-1055, 1071-1085, 1104-1113, 1121-1132, 1138-1150, 1187-1196, 1212-1221, 1227-1236, 1257-1262, 1264-1278, 1282-1294, 1307-1318, 1353-1370, 1382-1388, 1396-1409, 1434-1440, 1446-1454, 1465-1478, 1485-1513, 1516-1529, 1540-1545, 1563-1568, 1575-1593, 1607-1616, 1628-1645, 1648-1661, 1676-1682, 1689-1697, 1713-1719, 1739-1749, 1753-1758, 1763-1774, 1797-1803, 1807-1846, 1855-1874, 1877-1891, 1893-1907, 1912-1925, 1931-1943, 1955-1965, 1976-1990, 2032-2043, 2045-2051, 2099-2105, 2131-2138, 2161-2179, 2188-2199, 2205-2216, 2219-2227, 2235-2245, 2247-2267, 2277-2288, 2294-2304, 2314-2326, 2346-2358, 2365-2377, 2383-2402, 2407-2423, 2437-2450, 2454-2473, 2489-2497, 2525-2531, 2557-2570, 2580-2587, 2589-2599, 2621-2641, 2647-2653, 2661-2677, 2685-2690, 2697-2717, 2722-2733, 2739-2777, 2786-2793, 2801-2808, 2811-2822, 2825-2835, 2838-2845, 2859-2871, 2877-2883, 213-344, 954-1080 and 2524-2733 of Seq ID No 279; 10-16, 18-23, 28-41, 63-69, 77-91, 101-109, 118136, 146-153, 155-162, 168-179, 192-207, 217-226, 229-235, 239-254, 279-286, 294-307, 313-319, 334-341, 344-353, 363-377, 390-396 and 178-328 of Seq ID No 280; 18-42, 68-84, 89-95, 100-105, 107-115, 125-135, 154-177, 189-195, 205-228, 236-243, 252-259, 279-300, 309-316, 323-331, 340-351, 353-364, 377-402 and 85-97 of Seq ID No 281; 4-18, 26-32, 66-76, 100-126, 151-159, 178-186, 188-194, 200-210, 241-248, 253-259, 262-279, 284-291, 307-313, 315-322, 327-337, 376-386, 399-407, 432-441, 467-473, 487-497, 499-505, 543-549, 560-568, 585-593, 598-604, 608-614, 630-642, 647-653, 690-703, 717-730, 21-200 and 468-480 of Seq ID No 282; 17-49, 52-58, 62-73, 78-97, 100-117, 122-172, 185-190, 193-217, 225-236 and 33-42 of Seq ID No 283; 7-39, 50-58, 73-89, 96-107, 109-120, 126-142, 152-170, 178-202, 205-211, 224-244, 249-259, 261-270, 300-310, 312-325 and 158-169 of Seq ID No 284; 4-31, 40-64, 71-82, 85-92, 102-124, 126-139, 147-152, 159-173, 176-188, 195-207, 210-216, 234-241, 249-256, 258-276, 279-293, 296-302, 310-315, 349-356, 363-378, 380-403, 411-426, 435-441, 448-459, 463-476, 488-494 and 201-221 of Seq ID No 285; 5-13, 15-74, 87-104, 107-120, 123-129, 136-145, 150-191, 193-206, 227-248, 250-264, 278-302, 304-323, 332-378, 384-407, 409-419, 425-457, 462-471, 474-497, 511-545, 555-564, 571-578, 585-598, 640-647, 669-675, 682-691, 693-705, 729-743, 752-761, 772-780, 786-804, 808-818, 822-846, 858-880, 884-900, 910-939, 941-947, 962-971, 973-988, 998-1003, 1007-1027 and 236-259 of Seq ID No 286; 4-19, 27-68, 81-111, 121-160 and 60-79 of Seq ID No 287; 4-37, 40-46, 52-57, 199-205, 222-229, 236-244, 250-267, 269-282 and 27-197 of Seq ID No 288; 4-16, 24-30, 32-38, 63-75, 86-92, 98-111, 113-126, 160-165, 170-180, 198-204, 227-233, 239-245, 253-273, 308-314, 352-365, 382-387, 395-403, 423-429, 472-482, 484-493, 501-507, 518-526, 536-541, 543-550, 556-562, 586-600, 626-633, 649-661, 680-688 and 546-559 of Seq ID No 289; 16-33, 48-59, 63-71, 77-92, 94-109, 117-124, 139-151, 169-181, 184-227, 233-249, 251-261, 263-275, 282-294, 297-321, 326-332, 341-355, 383-399 and 258-272 of Seq ID No 290; 11-26, 31-39, 43-52, 55-62, 64-70, 80-94, 123-133, 135-141, 172-181, 185-206, 209-218, 224-230, 238-244, 251-262, 264-271, 290-301, 306-324, 333-340, 350-357, 367-375, 390-397, 434-441, 443-448, 77-226 and 350-429 of Seq ID No 291; 4-13, 22-27, 31-45, 50-59, 72-96, 99-114, 131-141, 143-150, 159-176, 180-186, 189-198, 208-214, 234-253, 271-287, 294-299, 310-366, 382-390, 398-416, 424-443 and 283-305 of Seq ID No 292; 9-26, 30-53, 62-72, 86-95, 112-122, 136-145, 153-160, 209-221, 227-237, 241-268, 281-288, 291-298, 308-314, 321-328, 336-346, 351-379, 388-397, 409-416, 423-433, 443-481, 511-519 and 213-232 of Seq ID No 293; 12-18, 25-31, 38-50, 59-67, 71-82, 96-126 and 76-88 of Seq ID No 294; 4-25, 39-44, 64-71, 74-88, 100-113, 128-138, 151-162, 164-177, 185-190, 204-213, 233-239, 246-254, 281-286, 293-306, 309-318, 333-347, 349-359, 385-398, 404-423, 458-465, 477-484, 490-499, 501-533, 554-566, 582-590, 596-616, 624-629, 631-639, 654-680, 694-720, 735-743 and 2-100 of Seq ID No 295; 4-16, 36-41, 52-75, 98-107, 109-117, 122-128, 133-139, 141-155, 159-165, 169-182, 187-193, 195-201, 211-224, 230-236, 247-269, 278-290 and 75-92 of Seq ID No 296; 7-21, 25-33, 37-43, 87-94, 103-120, 131-147, 168-174, 197-203, 207-212, 227-237, 247-257, 263-271, 279-287, 298-306, 320-325, 332-340, 363-374, 379-384, 390-401, 403-414, 428-433, 448-457, 462-475, 483-490, 513-519, 525-535, 543-554, 559-566, 571-620, 625-631, 636-642, 659-670, 688-706, 708-723, 770-779, 787-793, 796-807, 820-840, 848-854, 863-874, 895-905, 912-919, 934-942, 968-975, 983-1000, 1012-1019, 1026-1036, 1050-1060, 1064-1070, 1081-1091, 1094-1108, 1112-1118, 1140-1152, 1164-1169, 1172-1180, 1187-1192 and 732-748 of Seq ID No 297; 23-40, 42-59, 66-73, 78-97, 111-128, 130-141, 157-166, 178-183 and 53-71 of Seq ID No 298; 4-27, 38-44, 47-57, 59-85, 99-106, 114-121, 154-166, 181-186, 193-198, 238-244, 253-262, 272-278, 287-299, 314-320, 338-350, 358-368, 382-388, 407-416, 433-446, 456-461, 463-473 and 86-195 of Seq ID No 299; 5-24, 38-59, 64-80, 87-99, 105-126, 134-142, 149-163, 165-179, 181-202, 205-220, 227-233, 243-250, 257-263 and 87-245 of Seq ID No 300; 5-32, 47-53, 66-79, 81-97, 115-151, 155-174, 183-188, 196-210, 215-226, 230-238, 253-258, 263-270, 276-282, 295-301, 304-325, 334-344, 360-390, 397-412, 425-432, 434-462, 478-494, 508-526, 539-564, 571-579, 347-371 and 375-386 of Seq ID No 301; 4-15, 36-44, 49-56, 60-66, 68-82, 84-103, 109-115, 118-141, 147-154, 160-168, 176-185 and 26-39 of Seq ID No 302; 7-13, 23-33 and 13-21 of Seq ID No 303; 2-10 of Seq ID No 304; 4-9, 12-18, 35-42, 49-62 and 6-18 of Seq ID No 305; 19-25 and 1-13 of Seq ID No 306; 15-21, 27-45 and 12-25 of Seq ID No 307; 14-20 and 1-14 of Seq ID No 308; 4-18 and 13-26 of Seq ID No 309; 8-21 and 2-20 of Seq ID No 310; 4-14 and 4-16 of Seq ID No 311; 3-12 of Seq ID No 312; 6-14, 6-25, 35-57 and 2-14 of Seq ID No 313; 6-25, 35-57 and 17-31 of Seg ID No 314; 14-25, 32-46 and 5-19 of Seg ID No 315; 18-31 and 5-16 of Seg ID No 316; 19-24 and 4-26 of Seq ID No 317; 13-21, 29-34, 47-58, 61-73 and 36-47 of Seq ID No 318; 4-15 and 5-24 of Seq ID No 319; 6-18 of Seq ID No 320; 13-20 and 4-13 of Seq ID No 321; 15-23 of Seq ID No 322; 4-9 and 7-21 of Seq ID No 323; 1-10 of Seq ID No 324; 4-14 of Seq ID No 325; 4-17, 35-41, 46-89, 93-98 and 70-88 of Seq ID No 326; 1-13 of Seq ID No 327; 4-16, 26-32 and 25-38 of Seq ID No 328; 8-15, 23-28 and 4-17 of Seq ID No 329; 4-12 and 1-15 of Seq ID No 330; 4-29, 31-42, 52-58 and 6-16 of Seq ID No 331; 4-9, 24-32 and 9-19 of Seq ID No 332; 4-12, 18-27 and 5-18 of Seq ID No 333; 4-11, 37-56, 58-92 and 18-29 of Seq ID No 334; 8-28 and 20-35 of Seq ID No 335; 4-15 of Seq ID No 336; 4-23, 27-39, 55-63 and 35-58 of Seq ID No 337; 6-26, 28-54 and 28-47 of Seq ID No 338; 4-10, 38-52, 58-82 and 30-49 of Seq ID No 339; 4-22, 29-35, 44-50, 53-68, 70-80 and 20-33 of Seq ID No 340; 22-28, 30-36 and 18-33 of Seq ID No 341; 4-11, 13-21, 25-30 and 20-30 of Seq ID No 342; 10-22 and 10-23 of Seq ID No 343; 4-11 and 9-20 of Seq ID No 344; 14-25, 32-46 and 6-19 of Seq ID No 345; 5-30 and 14-33 of Seq ID No 346; 4-15, 28-35, 46-55, 59-65, 76-84 and 9-24 of Seq ID No 347; 27-33 and 5-19 of Seq ID No 348; 5-13 and 8-18 of Seq ID No 349; 9-22, 24-34 and 21-40 of Seq ID No 350; 4-17, 35-41, 46-89, 93-98 and 71-89 of Seq ID No 351; 4-12, 14-24 and 2-17 of Seq ID No 352; 9-17 and 5-16 of Seq ID No 353; 7-41, 48-58, 63-75, 80-89 and 43-53 of Seq ID No 354; 4-22, 25-30 and 4-14 of Seq ID No 355; 4-55 and 18-33 of Seq ID No 356; 262-280 of Seq ID No 179; 131-146 of Seq ID No 186; 207-224 of Seq ID No 188; 27-50, 203-217 and 313-325 of Seq ID No 189; 110-129 of Seq ID No 192; 156-179, 174-197, 192-215, 210-233, 228-251 and 246-267 of Seq ID No 190; 377-400 of Seq ID No 196; 34-43, 234-257 and 350-367 of Seg ID No 203; 304-327 of Seg ID No 207; 25-48, 43-66 and 61-82 of Seq ID No 222; 398-421, 416-439, 434-457, 452-475, 470-493, 488-511, 506-529, 524-547, 621-644, 639-664, 707-730, 725-748, 743-766, 761-784, 779-802, 797-820, 984-1007, 1002-1025, 1020-1043, 1038-1061, 1056-1079, 1074-1097, 1092-1115, 1286-1309, 1304-1327, 1322-1345, 1340-1363, 1358-1381, 1376-1399, 1394-1417, 1412-1435, 1430-1453, 1448-1471, 1466-1489 and 1484-1507 of Seq ID No 226; 188-211, 206-229, 224-247, 242-265, 260-283 and 278-296 of Seq ID No 227; 56-79 and 122-132 of Seq ID No 229; 35-46 of Seq ID No 231; 178-201, 196-219, 214-237, 232-255, 250-273, 268-291, 379-402, 397-420, 415-438, 433-456, 451-474, 642-665, 660-683, 678-701, 696-719, 714-737, 732-755, 750-773, 768-791, 899-922, 917-940, 935-958, 1037-1060, 1055-1078, 1073-1096 and 1091-1114 of Seq ID No 232; 330-346 of Seq ID No 233; 571-594, 589-612, 607-630, 625-648, 643-666 and 661-684 of Seq ID No 242; 188-207 of Seq ID No 244; 61-84, 308-331, 326-349, 344-367, 362-385, 380-403 and 398-421 of Seq ID No 249; 79-98, 345-366, 844-867, 870-887 and 890-905 of Seq ID No 258; 94-109 of Seq ID No 268; 188-207 of Seq ID No 272; 290-306 of Seq ID No 276; 826-849 of Seq ID No 277; 228-252, 247-270, 265-288, 283-306, 301-324, 955-978, 973-996, 991-1014, 1009-1032, 1027-1050, 1045-1068, 2533-2556, 2551-2574, 2569-2592, 2587-2610, 2605-2628 and 2623-2646 of Seq ID No 279; 86-109 and 104-127 of Seq ID No 288; 546-560 of Seq ID No 289; 260-271 of Seq ID No 290; 106-129, 124-147, 142-165, 160-183, 178-201 and 375-398 of Seq ID No 291; 284-307 of Seq ID No 292; 362-385 of Seq ID No 301.

- 15. A process for producing a *H. pylori* hyperimmune serum reactive antigen or a fragment thereof according to any one of the claims 11 to 14 comprising expressing the nucleic acid molecule according to any one of claims 1 to 7.
- 16. A process for producing a cell, which expresses a *H. pylori* hyperimmune serum reactive antigen or a fragment thereof according to any one of the claims 11 to 14 comprising transforming or transfecting a suitable host cell with the vector according to claim 8 or claim 9.
- 17. A pharmaceutical composition, especially a vaccine, comprising a hyperimmune serum-reactive antigen or a fragment thereof, as defined in any one of claims 11 to 14 or a nucleic acid molecule according to any one of claims 1 to 7.
- 18. A pharmaceutical composition, especially a vaccine, according to claim 17, characterized in that it further comprises an immunostimulatory substance, preferably selected from the group comprising polycationic polymers, especially polycationic peptides, immunostimulatory deoxynucleotides (ODNs), peptides containing at least two LysLeuLys motifs, neuroactive compounds, especially human growth hormone, alumn, Freund's complete or incomplete

- adjuvants or combinations thereof.
- 19. Use of a nucleic acid molecule according to any one of claims 1 to 7 or a hyperimmune serum-reactive antigen or fragment thereof according to any one of claims 11 to 14 for the manufacture of a pharmaceutical preparation, especially for the manufacture of a vaccine against *H. pylori* infection.
- 20. An antibody, or at least an effective part thereof, which binds at least to a selective part of the hyperimmune serum-reactive antigen or a fragment thereof according to any one of claims 11 to 14.
- 21. An antibody according to claim 20, wherein the antibody is a monoclonal antibody.
- 22. An antibody according to claim 20 or 21, wherein said effective part comprises Fab fragments.
- 23. An antibody according to any one of claims 20 to 22, wherein the antibody is a chimeric antibody.
- 24. An antibody according to any one of claims 20 to 23, wherein the antibody is a humanized antibody.
- 25. A hybridoma cell line, which produces an antibody according to any one of claims 20 to 24.
- 26. A method for producing an antibody according to claim 20, characterized by the following steps:
  - initiating an immune response in a non-human animal by administrating an hyperimmune serum-reactive antigen or a fragment thereof, as defined in any one of the claims 11 to 14, to said animal,
  - removing an antibody containing body fluid from said animal, and
  - producing the antibody by subjecting said antibody containing body fluid to further purification steps.
- 27. Method for producing an antibody according to claim 21, characterized by the following steps:
  - initiating an immune response in a non-human animal by administrating an hyperimmune serum-reactive antigen or a fragment thereof, as defined in any one of the claims 12 to 15, to said animal,
  - removing the spleen or spleen cells from said animal,
  - producing hybridoma cells of said spleen or spleen cells,
  - selecting and cloning hybridoma cells specific for said hyperimmune serum-reactive antigens or a fragment thereof,
  - producing the antibody by cultivation of said cloned hybridoma cells and optionally further purification steps.
- 28. Use of the antibodies according to any one of claims 20 to 24 for the preparation of a medicament for treating or preventing *H. pylori* infections.
- 29. An antagonist which binds to the hyperimmune serum-reactive antigen or a fragment thereof according to any one of claims 11 to 14.
- 30. A method for identifying an antagonist capable of binding to the hyperimmune serum-reactive antigen or fragment thereof according to any one of claims 11 to 14 comprising:
  - a) contacting an isolated or immobilized hyperimmune serum-reactive antigen or a fragment thereof according to any one of claims 11 to 14 with a candidate antagonist under conditions to permit binding of said candidate antagonist to said hyperimmune serum-reactive antigen or

- fragment, in the presence of a component capable of providing a detectable signal in response to the binding of the candidate antagonist to said hyperimmune serum reactive antigen or fragment thereof; and
- b) detecting the presence or absence of a signal generated in response to the binding of the antagonist to the hyperimmune serum reactive antigen or the fragment thereof.
- 31. A method for identifying an antagonist capable of reducing or inhibiting the interaction activity of a hyperimmune serum-reactive antigen or a fragment thereof according to any one of claims 11 to 14 to its interaction partner comprising:
  - a) providing a hyperimmune serum reactive antigen or a hyperimmune fragment thereof according to any one of claims 11-14,
  - b) providing an interaction partner to said hyperimmune serum reactive antigen or a fragment thereof, especially an antibody according to any one of the claims 20 to 24,
  - c) allowing interaction of said hyperimmune serum reactive antigen or fragment thereof to said interaction partner to form a interaction complex,
  - d) providing a candidate antagonist,
  - e) allowing a competition reaction to occur between the candidate antagonist and the interaction complex,
  - f) determining whether the candidate antagonist inhibits or reduces the interaction activities of the hyperimmune serum reactive antigen or the fragment thereof with the interaction partner.
- 32. Use of any of the hyperimmune serum reactive antigen or fragment thereof according to any one of claims 11 to 14 for the isolation and/or purification and/or identification of an interaction partner of said hyperimmune serum reactive antigen or fragment thereof.
- 33. A process for in vitro diagnosing a disease related to expression of the hyperimmune serum-reactive antigen or a fragment thereof according to any one of claims 11 to 14 comprising determining the presence of a nucleic acid sequence encoding said hyperimmune serum reactive antigen and fragment according to any one of claims 1 to 7 or the presence of the hyperimmune serum reactive antigen or fragment thereof according to any one of claims 11-14.
- 34. A process for *in vitro* diagnosis of a bacterial infection, especially a *H. pylori* infection, comprising analysing for the presence of a nucleic acid sequence encoding said hyperimmune serum reactive antigen and fragment according to any one of claims 1 to 7 or the presence of the hyperimmune serum reactive antigen or fragment thereof according to any one of claims 11 to 14.
- 35. Use of the hyperimmune serum reactive antigen or fragment thereof according to any one of claims 11 to 14 for the generation of a peptide binding to said hyperimmune serum reactive antigen or fragment thereof, wherein the peptide is selected from the group comprising anticalines.
- 36. Use of the hyperimmune serum-reactive antigen or fragment thereof according to any one of claims 11 to 14 for the manufacture of a functional nucleic acid, wherein the functional nucleic acid is selected from the group comprising aptamers and spiegelmers.
- 37. Use of a nucleic acid molecule according to any one of claims 11 to 14 for the manufacture of a functional ribonucleic acid, wherein the functional ribonucleic acid is selected from the group comprising ribozymes, antisense nucleic acids and siRNA.